Endoscopic pancreatic and biliary manometry in pancreatic, biliary, and papillary disease, and after endoscopic sphincterotomy and surgical sphincteroplasty

J A GREGG AND D L CARR-LOCKE

From the New England Baptist Hospital, Boston, Mass, USA and Leicester Royal Infirmary, Leicester

SUMMARY Endoscopic manometry was used to measure pancreatic duct, common bile duct, pancreatic duct sphincter and bile duct sphincter pressures in 43 healthy volunteers and 162 patients with a variety of papillary, pancreatic and biliary disorders. Common bile duct pressure was significantly raised after cholecystectomy, with common bile duct stones and papillary stenosis but pancreatic duct pressure only in papillary stenosis. After endoscopic sphincterotomy mean common bile duct pressure fell from 11.2 to 1.1 mmHg and pancreatic duct pressure from 18.0 to 11.2 mmHg. Distinct pancreatic duct sphincter and bile duct sphincter zones were identified as phasic pressures of 3–12 waves/minute on pull-through from pancreatic duct and common bile duct to duodenum. Pancreatic duct sphincter pressures were higher with common bile duct stones and stenosis whereas bile duct sphincter pressures were higher in pancreatitis and stenosis. Bile duct sphincter activity was present in 60% of patients after surgical sphincteroplasty but 21% of patients after endoscopic sphincterotomy. Endoscopic manometry facilitated the diagnosis of papillary stenosis, has allowed study of papillary pathophysiology and has shown a functional inter-relationship between the two sphincteric zones.

The role of the sphincter of Oddi in the pathogenesis of pancreatic and biliary diseases and their effects upon it have received little attention in human studies. Various pathophysiological mechanisms have been suggested to implicate dysfunction of the pancreatic-biliary sphincter apparatus in gall stone formation,1 2 pancreatitis3 4 and the postcholecystectomy syndrome5–7 but direct study of this area became possible only recently with the advent of endoscopic manometry.8–12 Previously the sphincter of Oddi was accessible only indirectly for manometric study by the use of intra-operative or postoperative bile duct pressure-flow systems.13 The diagnosis of papillary stenosis remains controversial despite an increasing literature on the subject since the 1950s.5 7 14–18 Radiological criteria based on ERCP studies19–21 have helped delineate this entity further but endoscopic manometric measurements10 11 21 22 would seem a promising method for accurate diagnosis.

Surgical and endoscopic operations on the sphincter of Oddi are frequently used in the management of choledocholithiasis and papillary stenosis.18 23–29 Little information is available on the effects of these procedures on the biliary-pancreatic sphincter apparatus and recent studies using endoscopic manometry10–12 30–32 have concentrated on measurements of common bile duct pressure10–12 30–32 and bile duct sphincter zone phasic activity.30–32 Numbers of patients studied before and after endoscopic sphincterotomy have been small and only one study reports changes in pancreatic duct pressure31 but not pancreatic duct sphincter zone dynamics.

We have previously reported our results of endoscopic manometry in healthy volunteers33 and now report our findings in a group of patients asymptomatic after cholecystectomy and those with common bile duct stones, pancreatitis, suspected papillary stenosis, and in patients after endoscopic sphincterotomy and surgical sphincteroplasty.
Methods

Patients
One hundred and sixty two patients gave written consent to undergo endoscopic manometry and their results were compared with 43 healthy volunteers (Group N) amalgamated from previous studies.35-36 These were 19 men and 24 women, aged 19 to 51 years, selected on the basis of an absence of a history of biliary, pancreatic or any other gastrointestinal or other disease, pregnancy, use of any regular medication or an alcohol consumption greater than 30 grams per day. The protocol was approved by the New England Baptist Hospital clinical investigation committee. Patients were divided into eight groups. (1) Group PC consisted of four patients, aged 25 to 75 years, asymptomatic after cholecystectomy in whom no papillary, biliary or pancreatic disorders were suspected or found. (2) Group CDS consisted of 13 patients, aged 43 to 86 years, with common bile duct stones of whom 10 had had a previous cholecystectomy. In none was there clinical, biochemical or radiological evidence of coexistent pancreatic disease. (3) Group P consisted of 12 patients, aged 23 to 75 years with acute or chronic relapsing pancreatitis documented on the basis of clinical, biochemical, ultrasonographic, radiological and secretory tests. In seven, pancreatitis was alcohol related and six had had a previous cholecystectomy. None was found to have an abnormal papilla during endoscopic cannulation and none had biliary calculus at the time of examination. (4) Group S consisted of 44 patients with a history of recurrent or constant biliary or pancreatic type pain considered to have papillary stenosis. This diagnosis was based on calibration of the papilla using a standard 1.7 mm diameter ERCP cannula, dilatation of the common bile duct of greater than 10 mm in the postcholecystectomy state and a pancreatic duct diameter of 5 mm or greater in the head of the pancreas with delayed drainage beyond 30 minutes of contrast medium from the ductal system after ERCP. Thirty eight patients in this group had had a previous cholecystectomy. (5) Group SPD consisted of five patients found to have stenosis of the pancreatic duct orifice at ERCP of whom four had had a previous surgical choledochal sphincteroplasty and one endoscopic sphincterotomy. Three patients had also had a previous failed distal pancreatic drainage procedure. (6) Group PSS consisted of 20 patients who had had a previous transduodenal surgical sphincteroplasty of the choledochal sphincter after cholecystectomy in whom symptoms had continued or returned suggesting pancreatic and/or biliary pain and had led to referral for ERCP. None had common bile duct stones. (7) Group PES consisted of 56 patients studied after endoscopic sphincterotomy of the choledochal sphincter performed for choledocholithiasis or papillary stenosis. (8) Group ESPD consisted of eight patients on whom endoscopic sphincterotomy of the pancreatic duct orifice was performed for stenosis. Six had had a previous surgical choledochal sphincteroplasty, two an endoscopic choledochal sphincterotomy and three had had a failed distal pancreatic drainage operation. All had severe episodic or continuous pancreatic-type pain before endoscopic sphincterotomy.

Endoscopic manometry was performed at the same session as, but immediately before ERCP as previously described.33 After an eight hour fast the subject lay comfortably in the left lateral or semiprone position after local pharyngeal anaesthesia had been administered by benzocaine spray. Sedation was then induced with intravenous diazepam and no other drugs were given until manometry recordings had been completed. Dudoenoscopy was performed with a Fujinon DUOX duodenoscope, and the manometry catheter, a modified Fujinon ERCP catheter with a 1-2 mm diameter side hole 4 mm from its sealed tip, was passed through the instrument channel into the duodenum with the transducer level with the subject's abdomen. Using a perfusion rate of 0-62 ml/min of 0-9% saline delivered by a Harvard 2681 infusion pump, an initial duodenal pressure was recorded. The papilla was then cannulated and the catheter deeply inserted into a duct. Aspiration of bile or pancreatic juice together with fluoroscopic verification allowed identification of the duct cannulated. Ductal pressures were then measured and the catheter then slowly withdrawn until a phasic high pressure zone was located on pull-through. A station recording was made and the distance of this zone from the papillary orifice could be read from the catheter markings. Duodenal pressure was again recorded after each pull-through manoeuvre as this was used as a zero reference for each recording. Ductal and duodenal pressures were read directly from each tracing and a mean of the peak and trough phasic pressures calculated from the phasic zone activity as previously described.33 Wave amplitude, frequency and duration were also calculated and groups were compared statistically using the Student's t test for unpaired data.

Results
The results of ductal pressure measurements are shown in Tables 1 and 2. Pancreatic duct (PD) pressure was not significantly different from normal
in asymptomatic postcholecystectomy subjects (PC), patients with common duct stones (CDS) and patients with pancreatitis (P) but was raised in papillary stenosis (S), stenosis of the pancreatic duct orifice (SPD) and in patients after surgical sphincteroplasty (PSS). Common bile duct (CBD) pressure was not significantly different from normal in Groups P and PSS but values in Groups PC, S and CDS were significantly raised. In nine patients (45%) of Group PSS there was no CBD-duodenal gradient present. Common bile duct pressure was also significantly raised in Groups CDS and S compared with asymptomatic postcholecystectomy subjects. Pancreatic duct pressure was higher than common bile duct pressure in all groups except those with common bile duct stones where the reverse was found.

The effect of endoscopic sphincterotomy is shown in Figures 1 and 2. Pancreatic duct pressure fell from 18.0±7.0 to 11.2±5.0 mmHg (p<0.001) and common bile duct pressure fell from 11.2±4.9 to 1-2±1.1 mmHg (p<0.001) with 39 subjects (70%) showing no common bile duct-duodenal pressure gradient. In the small group of patients undergoing sphincterotomy of the pancreatic duct orifice (ESPD) pancreatic duct pressure fell significantly from 22.4±7.6 to 8.0±6.0 mmHg (p<0.001).

Distinct zones of high phasic pressures with a mean frequency of six waves/min (range 3–12 waves/min) and a mean duration of seven seconds (6–2–8.4 seconds) were found on pull-through from pancreatic duct and common bile duct towards the duodenum at a distance of 4–5 mm from the papilla. These zones were designated pancreatic duct sphincter and bile duct sphincter zones respectively as we have previously discussed and examples are shown in Figure 3. Peak and trough pancreatic duct sphincter and bile duct sphincter pressures are shown in Tables 1 and 2. For pancreatic duct sphincter pressures there was no significant difference from normal in Groups PC, PES, and PSS but a significant rise was found in Groups P, CDS, S, and SPD. Significant falls in pancreatic duct sphincter pressures occurred after endoscopic sphincterotomy (Fig. 1) and in the eight subjects undergoing pancreatic duct orifice sphincterotomy (ESPD). Two subsequently showed no phasic activity and in the remaining six there were significant falls in pancreatic duct sphincter pressures (Fig. 1).

Table 1  Ductal and phasic sphincter pressures for normal subjects and patient groups

<table>
<thead>
<tr>
<th>Patient group</th>
<th>Pancreatic duct</th>
<th>Common bile duct</th>
<th>Bile duct sphincter</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Peak</td>
<td>Trough</td>
<td>Peak</td>
</tr>
<tr>
<td>N (43)</td>
<td>10.7±3.9</td>
<td>15.9±6.6</td>
<td>51.2±6.7</td>
</tr>
<tr>
<td>PC (4)</td>
<td>14.0±4.0</td>
<td>11.0±7.1</td>
<td>6.0±1.8</td>
</tr>
<tr>
<td>CDS (13)</td>
<td>10.0±2.0</td>
<td>30.3±15.7</td>
<td>14.2±5.8</td>
</tr>
<tr>
<td>P (12)</td>
<td>15.4±7.9</td>
<td>20.1±10.9</td>
<td>7.1±6.5</td>
</tr>
<tr>
<td>S (44)</td>
<td>18.9±6.8‡</td>
<td>23.5±11.1‡</td>
<td>9.9±4.0†</td>
</tr>
</tbody>
</table>

Given in mmHg as mean ± 1SD. Significant differences from normal values are given by * p<0.01, † p<0.005, ‡ p<0.001.
N = normal. PC = postcholecystectomy. CDS = common duct stone. P = pancreatitis. S = papillary stenosis.

Table 2  Ductal and phasic sphincter pressures for normal subjects and patient groups

<table>
<thead>
<tr>
<th>Patient group</th>
<th>Pancreatic duct</th>
<th>Common bile duct</th>
<th>Bile duct sphincter</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Peak</td>
<td>Trough</td>
<td>Peak</td>
</tr>
<tr>
<td>N (43)</td>
<td>10.7±3.9</td>
<td>15.9±6.6</td>
<td>2.0±1.7</td>
</tr>
<tr>
<td>SPD (5)</td>
<td>22.4±10.9‡</td>
<td>25.2±9.2†</td>
<td>37.0±12.3‡(12)</td>
</tr>
<tr>
<td>PSS (20)</td>
<td>17.0±7.5‡</td>
<td>13.0±8.5</td>
<td>1.1±1.1(17)</td>
</tr>
<tr>
<td>PES (56)</td>
<td>11.2±5.0</td>
<td>16.9±8.5</td>
<td>31.2±9.8‡(12)</td>
</tr>
<tr>
<td>ESPD (8)</td>
<td>8.0±6.0</td>
<td>9.6±7.4§(6)</td>
<td>—</td>
</tr>
</tbody>
</table>

Given in mmHg as mean ± 1SD. Significant differences from normal values given by * p<0.01, † p<0.005, ‡ p<0.001.
Number of subjects in parentheses.
N = normal, SPD = pancreatic duct stenosis, PSS = surgical sphincteroplasty, PES = endoscopic sphincterotomy, ESPD = pancreatic sphincterotomy.
Fig. 1  Pancreatic duct and pancreatic duct sphincter pressures in 43 healthy volunteers (normal), 56 patients before and after endoscopic sphincterotomy and 8 patients before and after endoscopic sphincterotomy of the pancreatic duct orifice. Values are mean ± 1SD pancreatic duct pressure, phasic sphincter pressures. Statistical differences from pre-endoscopic sphincterotomy measurements are shown by *.

Peak and trough bile duct sphincter pressure results are shown in Tables 1 and 2. There was no significant difference from normal in Groups PC and CDS but a significant rise was found in Groups P and S. After surgical sphincteroplasty 12 of the 20 patients showed significantly lower pressures than normal and in the other eight of this group no detectable phasic activity was present. After endoscopic sphincterotomy no detectable bile duct sphincter activity was present in 44 of the 56 subjects and in the remaining 12 pressures were significantly lower than presphincterotomy values and normal subjects.

In patients with a diagnosis of papillary stenosis...
two patterns of pull-through recordings were obtained as shown in Figure 4. One was characterised by high amplitude waves with raised peak and trough pressures together with a raised ductal pressure and the other pattern exhibited a high pressure zone extending over 2-3 mm with little or no phasic activity at the expected site together with a raised ductal pressure.

Table 3 shows the percentage of patients in Groups CDS, P and S with ductal and phasic pressure values greater than 2SD above the normal mean. This revealed that 85% of patients with common bile duct stones and 75% with papillary stenosis had common bile duct pressures greater than 9.6 mmHg. In addition, 40% of patients with papillary stenosis had pancreatic duct pressures above the limit of 18.5 mmHg and two-thirds of this group had raised peak pancreatic duct sphincter and peak bile duct sphincter pressures above the normal limits. Table 4 expresses the same data as the percentage in each group with between one and six abnormal values based on the above mentioned limits. In patients with common bile duct stones the majority had only one abnormal value, in pancreatitis five patients (42%) had no abnormal value but in papillary stenosis the majority of patients had two or more abnormal values.

The differences between a ductal and trough phasic sphincter pressures were 7.3±4.9 and 12.0±5.7 mmHg in normal subjects for pull-through recordings from pancreatic duct and common bile duct respectively. Values in patient groups were not significantly different except those with papillary stenosis where this difference was significantly raised to 13.1±10.3 mmHg (p<0.02) and 18.6±13.3 mmHg (p<0.01) for pancreatic duct and common bile duct recordings respectively.

Discussion

Despite detailed anatomical descriptions of the muscle fibres associated with the terminal parts of the common bile duct and pancreatic duct in man and many other species over the last century there is still debate concerning the functional activity of these sphincteric structures in man and little is known of the pathophysiological changes which may take place in this area in patients with biliary and pancreatic disease. Endoscopic manometry has been used by a number of groups to investigate different disease states with varying results and some groups have also attempted to provide diagnostic criteria for papillary stenosis. We33-35 and others have shown common bile duct-duodenal and pancreatic duct-duodenal pressure gradients and phasic pressures on pull-through from common bile duct to duodenum which, we believe, represent components of the sphincter of Oddi. Further evidence comes from the results in this and other studies showing a fall in the common bile duct-

---

**Fig. 4** Endoscopic manometry recordings from (a) normal subject showing duodenal pressure (D) and bile duct pull-through phasic activity (bile duct sphincter) with small waves simultaneous with respiration (R), (b) patient with papillary stenosis after cholecystectomy showing raised common bile duct (CBD) and phasic pressures, (c) patient with papillary stenosis following cholecystectomy showing high plateau pressure replacing phasic activity in presumed stenotic segment (SS), and (d) patient shown in (c) after endoscopic sphincterotomy with loss of all phasic activity and CBD-duodenal pressure gradient.
Table 3  Percentage of patients in three disease groups

<table>
<thead>
<tr>
<th>Pressures</th>
<th>Group N, mean±2SD mmHg (43)</th>
<th>% greater than value in first column</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Group CDS (13)</td>
</tr>
<tr>
<td>Pancreatic duct</td>
<td>18-5</td>
<td>0</td>
</tr>
<tr>
<td>Pancreatic duct sphincter</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trough</td>
<td>29-2</td>
<td>15-4</td>
</tr>
<tr>
<td>Peak</td>
<td>63-9</td>
<td>15-4</td>
</tr>
<tr>
<td>Common bile duct</td>
<td>9-6*</td>
<td>85</td>
</tr>
<tr>
<td>Bile duct sphincter</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trough</td>
<td>25-8</td>
<td>7-7</td>
</tr>
<tr>
<td>Peak</td>
<td>64-5</td>
<td>23-1</td>
</tr>
</tbody>
</table>

(Number of patients in parentheses) with values above an arbitrary normal limit of mean ± 2SD from Normal.
* mean ± 2SD from Postcholecyst.

duodenal gradient to near zero\textsuperscript{10-12 30-32} and abolition of phasic activity\textsuperscript{30 32} after endoscopic sphincterotomy. We have previously discussed our contention that a separate manometrically measurable pancreatic duct sphincter zone exists on pull-through from pancreatic duct to duodenum\textsuperscript{33} which is still present after destruction of the choledochal sphincter. The falls in pancreatic duct and pancreatic duct sphincter pressures after sphincterotomy imply a functional interdependence between the bile duct sphincter and pancreatic duct sphincter zones as suggested by the anatomical inter-relationships of Boyden's sphincter choledochus and sphincter pancreaticus.\textsuperscript{37}

A consistent finding throughout studies in which pancreatic duct and common bile duct pressures have been measured in the same individuals\textsuperscript{10 22 31 40 44-46} is that pancreatic duct pressure is higher than common bile duct pressure and we have had similar results in studies of normal subjects.\textsuperscript{33-35} This may explain the mechanism underlying a recent report of pancreatic enzymes present in common bile duct bile.\textsuperscript{1} The difference was reversed in our patients with common bile duct stones, a finding not supported by others,\textsuperscript{31 40} and was due to the high common bile duct pressures compared with controls. This may be relevant to the mechanism of gall stone related acute pancreatitis\textsuperscript{3} but the mechanism of high common bile duct pressures in this situation is not known.

Our findings in patients with pancreatitis show rises of ductal and sphincteric pressures and there was no difference in values between alcohol related and non-alcohol related cases. The relevance of these findings to pathogenetic mechanisms of pancreatitis or the effects of pancreatitis on sphincter function are unknown but as some patients have values outside the proposed normal limits (Table 3) there may be a degree of sphincter dysfunction and outflow obstruction in a proportion.

Table 4  Percentage of patients in three disease groups with none or 1–6 abnormal values based on the arbitrary limit shown in Table 1.

<table>
<thead>
<tr>
<th></th>
<th>Group CDS, % (13)</th>
<th>Group P, % (12)</th>
<th>Group S, % (44)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No abnormal</td>
<td>0</td>
<td>41-8</td>
<td>0</td>
</tr>
<tr>
<td>1 abnormal</td>
<td>69-2</td>
<td>16-6</td>
<td>18-2</td>
</tr>
<tr>
<td>2 abnormal</td>
<td>15-4</td>
<td>25</td>
<td>29-5</td>
</tr>
<tr>
<td>3 abnormal</td>
<td>7-7</td>
<td>8-3</td>
<td>27-3</td>
</tr>
<tr>
<td>4 abnormal</td>
<td>7-7</td>
<td>8-3</td>
<td>11-4</td>
</tr>
<tr>
<td>5 abnormal</td>
<td>0</td>
<td>0</td>
<td>6-8</td>
</tr>
<tr>
<td>6 abnormal</td>
<td>0</td>
<td>0</td>
<td>6-8</td>
</tr>
</tbody>
</table>

(Number of patients in each group in parentheses.)

Gregg and Carr-Locke
diagnosis of papillary stenosis. A functional inter-
relationship between the sphincteric zones of the
terminal bile duct and pancreatic duct is suggested
but the pancreatic duct sphincter can preserve its
independent activity after complete incision of the
choledochal sphincter.

This work was supported by grants from the
Katherine Gavriluk and Sara Jordan Funds, New
England Baptist Hospital, Boston, Massachusetts
and Dr Carr- Locke is in receipt of grants from the
Wellcome Research Travel Fund, London, the
Leicester Area Health Authority and the P and C
Hickinbotham Trust, Leicester, England. We are
grateful to Elizabeth Knight RN and Louise Jackson
RN who assisted during endoscopic manometry and
Fujinon Optical Inc. New York, USA for help with the
equipment.

References
1 Anderson MC, Hauman RL, Suriyapa C, Schiller WR.
Pancreatic enzyme levels in bile of patients with extra
hepatic biliary tract disease. Am J Surg 1979; 137:
301–6.
2 Hauman RL, Gramatica L, Anderson MC. Effect of
specific pancreatic enzymes on the gallbladder. Surg
3 White TT. The part that the sphincter of Oddi plays in
the etiology of pancreatitis. In: The sphincter of Oddi.
Proc 3rd Gastroenterol Symp. Nice 1976, Basel:
4 Mouiel J, Bourgeon R, Chauvin P, Bertrand JC,
Giaume F, Rey JF. Pancreatitis due to obstruction of
Oddi’s sphincter. In: The sphincter of Oddi. Proc 3rd
163–74.
5 Tondelli P, Gyr K, Stalder GA, Allgower M. The post
cholecystectomy syndrome. Clin Gastroenterol 1979;
8: 487–505.
6 Schein CJ. Post cholecystectomy syndromes. A clinical
approach to etiology, diagnosis and management. New
7 Delmont J. An attempt to collate. In: The sphincter of
8 Vondrasek P, Eberhard G, Classen M. Endoscopic
9 Nebel OT. Manometric evaluation of the papilla of
10 Rosch W, Koch H, Demling L. Manometric studies
during ERCP and endoscopic papillotomy. Endoscopy
1976; 8: 30–3.
11 Hagenmuller F, Ossenberg FW, Classen M. Duodenosco-
nic manometry of the common bile duct. In: The
sphincter of Oddi. Proc 3rd Gastroenterol Symp. Nice
12 Geenen JE, Hogan WJ, Shaffer RD, Stewart ET,
Dodds WJ, Arndorfer RC. Endoscopic electrosur-
gical papillotomy and manometry in biliary tract
13 Wong HN, Frey CF, Gagic NM. Intraoperative
common duct pressure and flow measurements. Am J
14 Cattell RB, Colcock BP. Fibrosis of the sphincter of
15 Acosta JM, Civantos F, Nardi GL, Castleman B.
Fibrosis of the papilla of Vater. Surg Gynecol Obstet
16 Youngneaux JP, Bauwens E, Van Outryve L,
Yvergneaux E. Benign stenosis of the papilla of Vater.
17 Gregg JA, Clark G, Barr C, McCartney A, Milano A,
Volcjak C. Post cholecystectomy syndrome and its
139: 374–8.
18 Siegel HJ. Endoscopic management of choledocholithiasis
and papillary stenosis. Surg Gynecol Obstet 1979;
19 Zimmon DS, Ferrara TP and Clemettis AR. Radiology of
papilla of Vater stenosis. Gastrointest Radiol 1978;
20 Anacker H, Weiss HD and Kramann B. Endoscopic
retrograde pancreatico-cholangiography in chronic
diseases of the pancreas and papillary stenosis.
21 Delmont J. An attempt to collate. In: The sphincter of
Oddi. Proceedings of the Third Gastrointestinal
22 Bar-Meir S, Geenen JE, Hogan WJ, Dodds WJ,
Stewart ET, Arndorfer RC. Biliary and pancreatic duct
pressures measured by ERCP manometry in patients
with suspected papillary stenosis. Dig Dis Sci 1979;
23 Geenen JE, Vennes JA, Silvis SE. Resume of a
seminar on endoscopic retrograde sphincterotomy.
24 Cotton PB. Non-operative removal of bile duct stones
67: 1–5.
25 Nakajima M, Kizu M, Aksaka Y and Kawai K. Five
years experience of endoscopic sphincterotomy in
26 Reiter JJ, Bayer HP, Men Nicken C, Manigold BC.
Results of endoscopic papillotomy. A collective
experience from nine endoscopic centres in West
27 Safrany L. Endoscopic treatment of biliary tract
28 Classen M, Ossenberg FW. Non-surgical removal of
29 Jones SA. Sphincteroplasty (not sphincterotomy) in
the treatment of biliary tract disease. Surg Clin N Am
30 Funch-Jensen P, Csendes A, Kruse A, Oster MJ,
Amdrup E. Common bile duct and Oddi sphincter
pressure before and after endoscopic papillotomy in
patients with common bile duct stones. Ann Surg 1979;
190: 176–8.


